

**REMARKS**

Reconsideration of this application is respectfully requested.

Claims 7, 27, and 28 have been canceled. Claims 8-12 have been amended.

Upon amendment, claims 6, 8-12, and 25-26 are pending in the application.

Support for amended claims 8-12 can be found throughout the specification, for example, in original claims 8-12.

Applicants have also amended the specification at page 4, line 14 and page 15, line 7 to replace "∇-helices" with "α-helices." Support for this amendment can be found in the reference entitled Skouloubris et al. at page 4520 in the legend to Figure 3 (*Infection and Immunity*, 66(9): 4517-4521 (1998)). Skouloubris et al. is incorporated into the specification at page 15, line 27, was submitted to the Office, and was initialed by the Examiner on the PTO Form 1449 in the Office Action mailed April 27, 2004. Figure 3 of Skouloubris et al. reports the same graphic as shown at Figures 3(A) and 3(B) of this application, and indicates that the boxed regions show predicted transmembrane α-helices. Applicants further submit that one of skill in the art would recognize that α-helices denote a common and well characterized protein domain, whereas ∇-helices likely arose from a typographical error. A courtesy copy of the Skouloubris et al. reference is provided at Tab B.

Applicants submit that these amendments are fully supported by the specification, do not introduce new matter or require a further search of the art, and respectfully request their entry.

## **Priority**

The Office contends that Applicants have failed to comply with one or more conditions for receiving the benefit of an earlier filing date under 35 U.S.C. § 120. (Office Action at item 1.) In part, the Office contends that the first sentence of the specification contains errors in the filing dates of Application Serial Nos. 09/742,361 and 09/107,383. Applicants have amended the first sentence of the specification to contain the correct filing dates.

The Office also contends that the issue date for U.S. Patent No. 5,190,667 should be set forth in the first sentence of the Specification. In an effort to facilitate prosecution, Applicants have amended the first sentence to set forth the issue date.

The Office also asserts that PCT/EP99/04490 does not claim priority to U.S. Application No. 09/107,383, and therefore is not a Continuation-in-Part of U.S. Application No. 09/107,383. (Office Action at item 1d.) Applicants respectfully traverse. As proof that the stated priority claim is correct, Applicants direct the Office's attention to the Priority Data and Related Application Data listed on the cover page of WO 00/00634, the International Publication of Application No. PCT/EP99/04490 (attached at Tab A). Applicants respectfully submit (1) that the Priority Data for PCT/EP99/04490 lists U.S. Application No. 09/107,383, and (2) that PCT/EP99/04490 is Related by Continuation-in-Part to U.S. Application Serial No. 09/107,383.

Given the correction of the filing dates, the inclusion of the issue date, and the evidence that the stated priority claim is correct, Applicants respectfully request the benefit of an earlier filing date under 35 U.S.C. § 120.

## Drawings

The Office objected to drawings 3A and 3B because the Brief Description of the Drawings did not refer to each of the frames shown in the figures. (Office Action at Item 3.) Applicants have Amended the Brief Description of the Drawings to refer to both Figure 3A and to Figure 3B, and respectfully request withdrawal of this objection.

## Claim Rejections under 35 U.S.C. § 102

*Labigne et al.* (U.S. Patent No. 6,258,359)

The Office rejected claims 6-7, 9-11, and 25-27 under 35 U.S.C. § 102(e) as being anticipated by *Labigne et al.* (U.S. Patent No. 6,258,359). (Office Action at item 5.) In particular, the Office alleges that *Labigne et al.* disclose a method of treating *H. pylori* infection through formulating Urel polypeptide into a composition to induce antibodies that interfere with the activation process of the urease apoenzyme, wherein the antibodies are induced *in vivo* in man or animal. (Office Action at item 5.) Claims 7 and 27 have been canceled, so the rejection of these claims is now moot. Applicants respectfully traverse the rejection of claim 6, amended claims 9-11, and amended claims 25 and 26.

*Labigne et al.* state that antibodies raised against Urel gene product preferentially are capable of interfering “with the activation process of the urease apoenzyme.” (*Labigne et al.*, col. 10, ll. 24-27.) *Labigne et al.* also state that antibodies “may be used in therapeutic treatment of *Helicobacter pylori* infection in man by blocking the urease maturation process.” (*Labigne et al.*, col. 11, ll. 30-32.) However, the Urel deficient *H. pylori* strain used in the method of the instant invention is urease

positive, and thus is not identified by its capacity to block the urease maturation process.

Applicants further point out that in order to properly establish that Labigne et al. anticipates Applicants' claimed invention under 35 U.S.C. § 102(e), each and every element of the claim in issue must be found, either expressly described or under principles of inherency, in that single reference. Furthermore, "[t]he identical invention must be shown in as complete detail as is contained in the . . . claim." See M.P.E.P. §2131, 8th Ed., Aug. 2001, p. 2100-69, quoting *Richardson v. Suzuki Motor Co.*, 868 F.2d 1126, 1236, 9 U.S.P.Q.2d 1913, 1920 (Fed. Cir. 1989). Finally, "[t]he elements must be arranged as required by the claim." M.P.E.P. §2131, p. 2100-69. Regarding the 35 U.S.C. § 102(e) rejection, Labigne et al. does not teach each and every element of Applicants' present invention as claimed.

Claim 6 recites, among other things, "testing and comparing the response to extracellular pH and the sensitivity to acidity of the parental strain to a strain deficient in Urel and/or of a Urel deficient strain complemented with a plasmid carrying *urel* in the presence or absence of said active molecule." Labigne et al. nowhere discloses testing and comparing the response to extracellular pH and the sensitivity to acidity. Further, Labigne et al. nowhere discloses the selective use of Urel parental and deficient strains. Because Labigne does not teach each and every element of claim 6, Labigne et al. cannot anticipate claim 6, or dependent claims 9-11, 25, and 26.

Labigne et al. further fails to teach elements of dependent claims 9-11, 25, and 26. For example, claim 9 recites a molecule that inactivates Urel by inhibiting its properties in *H. pylori* resistance to acidity. Labigne et al. does not disclose whether or

not Urel antibodies affect the resistance to acidity of *H. pylori*. Claim 10 recites a molecule that inactivates Urel by inhibiting its property as a transporter. Labigne et al. does not address whether or not its antibodies inhibit the transport properties of Urel. Claim 11 recites a molecule that inactivates Urel “by inhibiting an interaction between Urel and other *H. pylori* proteins.” Labigne et al. does not disclose whether or not Urel antibodies inhibit an interaction between Urel and other *H. pylori* proteins. Claim 26 recites a molecule that “specifically inhibits Urel transporter properties either in ammonia export or in urea export or import. Labigne et al. does not address whether or not Urel antibodies specifically inhibit Urel transporter properties in ammonia export or in urea export or import. Because Labigne does not teach each and every element of claims 9-11, 25, and 26, Labigne et al. cannot anticipate these claims.

*Iversen et al. (U.S. Patent No. 6,124,271)*

The Office rejected claims 6-7, 12, and 28 under 35 U.S.C. § 102(e) as being anticipated by Iversen et al. (U.S. Patent No. 6,124,271). (Office Action at item 6.) In particular, the Office alleges that Iversen discloses administering *H. pylori* urease oligomer. Applicants have canceled claims 7 and 28, thus, the rejection of these claims is moot. Applicants respectfully traverse this rejection with respect to claim 6 and amended claim 12.

Claim 6 recites treating or preventing *H. pylori* infection with a molecule displaying a differential effect on parental and Urel deficient strains. Iversen et al. does not teach a molecule for treating *H. pylori* infection based on displaying a differential effect on parental and Urel deficient strains. A claim is anticipated only if each and

every element as set forth in the claim is found in a single prior art reference. M.P.E.P.

§ 2131.01 Because Iversen et al. does not teach selecting a molecule displaying a differential effect on parental and Urel deficient strains, it cannot anticipate claim 6 or dependent claim 12.

*Nakazawa et al. (U.S. Patent No. 5,214,053)*

The Office rejected claims 6-11 and 25-28 under 35 U.S.C. § 102(b) as being anticipated by Nakazawa et al. (U.S. Patent No. 5,214,053). (Office Action at items 7-8.) In particular, the Office alleged that Nakawaza et al. inherently anticipates the claimed invention, as Nakazawa et al. discloses thiourea compositions that have antimicrobial activity against *H. pylori*. (Office Action at item 8.) Applicants have canceled claims 7, 27, and 28, so the rejection of these claims is now moot. Applicants respectfully traverse with respect to claims 6, 8-11, 25, and 26.

As stated above, claim 6 recites treating or preventing *H. pylori* infection with a molecule displaying a differential effect on parental and Urel deficient strains. Nakazawa et al. does not teach the use of a molecule displaying a differential effect on parental and Urel deficient strains, and thus cannot anticipate claim 6. Claim 6 also recites comparing the response to extracellular pH and the sensitivity to acidity of a parental strain and a Urel deficient strain, which also is not disclosed in Nakazawa et al.

The Federal Circuit has explained that “[u]nder the doctrine of inherency, if an element is not expressly disclosed in a prior art reference, the reference will still be deemed to anticipate a subsequent claim if the missing element is necessarily present in the thing described in the reference, and that it would be so recognized by persons of

ordinary skill." *Rosco Inc v. Mirror Lite Co.*, 64 U.S.P.Q.2d 1676, 1680 (Fed. Cir. 2002) (emphasis added). Applicants submit that the thiourea compositions of Nakazawa et al. need not necessarily show differential effect in sensitivity to acidity between a parental *Helicobacter* and a strain deficient in Urel, as there is no teaching in Nakazawa et al. as to which subunit of the urease enzyme is disrupted by the thiourea compositions. Accordingly, the rejection of claim 6, as well as dependent claims 8-11 and 25-26, is improper.

*Hartmann (U.S. Patent No. 5,900,410)*

The Office also rejected claims 6-11 and 27 under 35 U.S.C. § 102(b) as being anticipated by Hartmann (U.S. Patent No. 5,900,410) as evidenced by Nawaz et al. (1994). (Office Action at item 9.) In particular, the Office contends that Hartmann discloses a method of treating *H. pylori* infection by administering a divalent cation of Mg, an antibiotic, and urea together with a pharmaceutically acceptable carrier. The Office alleges that Hartmann inherently anticipates the instantly claimed invention, as Nawaz et al. provides evidence that aliphatic amidases (such as Urel) are inhibited by divalent cations. (Office Action at item 9.) Applicants have cancelled claims 7 and 27, so the rejection is now moot with respect to these claims. Applicants respectfully traverse with respect to claims 6 and 8-11.

Nawaz et al. shows that Mg cation partially inhibits a purified aliphatic amidase from a *Rhodococcus* species. From this disclosure, the Office concludes that the administration of divalent cations in Hartmann can be used to treat infections of *H. pylori*. Applicants respectfully point out that "[i]nherency may not be established by

probabilities or possibilities. The mere fact that a certain thing may result from a given set of circumstances is not sufficient to establish inherency.” *Scaltech, Inc. v. Retec/Tetra, L.L.C.*, 51 U.S.P.Q.2d 1055, 1059 (Fed. Cir. 1999). Applicants submit that the Office has improperly concluded that Mg cation can be used to treat or prevent *H. pylori* infections in humans, as neither Hartmann et al. nor Nawaz et al. provide any indication of the *in vivo* effect of Mg cation on *H. pylori* Urel.

Applicants further submit that neither Hartman et al. nor Nawaz et al. provide any evidence that Mg divalent cation displays a differential effect on parental and Urel deficient strains, as is recited in claim 6. Under 35 U.S.C. § 102(b), each and every element of the claim in issue must be found, either expressly described or under principles of inherency, in that single reference. M.P.E.P. § 2131. Because neither Hartmann et al. nor Nawaz et al. explicitly or implicitly teach every element of claim 6, they cannot anticipate claim 6. Accordingly, the rejection of claim 6, as well as dependent claims 8-11 and 25-26, is improper.

#### WO 97/26908

The Office also rejected claims 6-7 under 35 U.S.C. § 102(a) as being anticipated by WO 97/26908. (Office Action at item 10.) In particular, the Office alleges that WO 97/26908 discloses a method of treating *Helicobacter pylori* infection comprising thiocyanate, as Gregoriou et al. provides evidence that thiocyanate is an inhibitor of amidase activity. (Office action at item 10.) Applicants respectfully traverse.

Gregoriou et al. shows that sodium cyanate inhibits purified amidase from *Pseudomonas aeruginosa* strain AI3. (Gregoriou et al., abstract.) From this disclosure,



the Office concludes that the thiocyanate in WO 97/26908 can be used to treat *H. pylori* infection. Applicants submit that the Office has improperly concluded that thiocyanate can be used to treat or prevent *H. pylori* infections in humans, as neither Hartmann et al. nor Nawaz et al. provide any indication of the *in vivo* effect of thiocyanate on *H. pylori* Urel.

Applicants further submit that neither WO97/26908 nor Gregorious et al. provide any evidence that thiocyanate displays a differential effect on parental and Urel deficient strains, as is recited in claim 6. Under 35 U.S.C. § 102(b), each and every element of the claim in issue must be found, either expressly described or under principles of inherency, in that single reference. M.P.E.P. § 2131. Because neither WO97/26908 nor Gregorious et al. explicitly or implicitly teach every element of claim 6, they cannot anticipate claim 6. Accordingly, the rejection of claim 6 is improper.

*Zopf (U.S. Patent No. 5,514,660)*

The Office asserts that claims 6 and 7 are rejected under 35 U.S.C. § 102(b) as being anticipated by Zopf (U.S. Patent No. 5,514,660) as evidenced by Gregoriou et al. (1979). In particular, the Office alleges that Zopf discloses a method of treating *H. pylori* infections by administering a cyanate derivative together with other antiulcerative compounds (see claims 10-11). Claim 7 has been canceled, so the rejection of this claim is now moot. Applicants respectfully traverse with respect to claim 6.

As stated above, claim 6 recites treating or preventing *H. pylori* infection with a molecule displaying a differential effect on parental and Urel deficient strains. Because

Zopf et al. does not teach a molecule displaying a differential effect on parental and Urel deficient strains, it cannot anticipate claim 6. See M.P.E.P. § 2131.01

Furthermore, as stated previously, Gregoriou et al. shows that sodium cyanate inhibits purified amidase from *Pseudomonas aeruginosa* strain AI3. (Gregoriou et al., abstract.) From this, the Office concludes that a cyanate derivative can be administered *in vivo* to inhibit *H. pylori* Urel. Applicants respectfully point out that “[i]nherency may not be established by probabilities or possibilities. The mere fact that a certain thing may result from a given set of circumstances is not sufficient to establish inherency.” *Scaltech, Inc. v. Retec/Tetra, L.L.C.*, 51 U.S.P.Q.2d 1055, 1059 (Fed. Cir. 1999). Accordingly, Applicants respectfully submit that the rejection of claim 6 under Zopf et al. is improper.

In view of the foregoing amendments and remarks, Applicants respectfully request the withdrawal of the rejections under 35 U.S.C. § 102.

## **Conclusion**

In view of the foregoing amendments and remarks, Applicants respectfully request reconsideration and reexamination of this application and the timely allowance of the pending claims.

Please grant any extensions of time required to enter this response and charge any additional required fees to our deposit account 06-0916.

Respectfully submitted,

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**Attachments:**

Cover page of PCT International Publication No. WO 00/00634  
Skouloubris et al. *Infection and Immunity*; 66(9): 4517-4521 (1998)

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## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

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(72) Inventors; and (75) Inventors/Applicants (for US only): DE REUSE, Hilde [BE/FR]; 49, rue Rouelle, F-75015 Paris (FR). SK-OULOUBRIS, Stéphane [FR/FR]; 45, rue de l'Amiral Mouchez, F-75013 Paris (FR). CUSSAC, Valérie [FR/FR]; 2 bis, impasse Morlet, F-75011 Paris (FR). LABIGNE, Agnès [FR/FR]; 47, avenue Beauséjour, F-91440 Bures-sur-Yvette (FR).			Published Without international search report and to be republished upon receipt of that report.
(54) Title: METHODS OF INHIBITING <i>HELICOBACTER PYLORI</i>			
(57) Abstract  This invention relates to methods of screening molecules capable of inhibiting the survival of <i>Helicobacter pylori</i> <i>in vivo</i> by specifically inhibiting the activity of UreI, to the molecules identified by these methods, and to the use of these molecules to treat or prevent <i>H. pylori</i> infection.			

## The *Helicobacter pylori* UreI Protein Is Not Involved in Urease Activity but Is Essential for Bacterial Survival In Vivo

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We produced defined isogenic *Helicobacter pylori* *ureI* mutants to investigate the function of UreI, the product of one of the genes of the urease cluster. The insertion of a *cat* cassette had a strong polar effect on the expression of the downstream urease genes, resulting in very weak urease activity. Urease activity, measured in vitro, was normal in a strain in which *ureI* was almost completely deleted and replaced with a nonpolar cassette. In contrast to previous reports, we thus found that the product of *ureI* was not necessary for the synthesis of active urease. Experiments with the mouse-adapted *H. pylori* SS1 strain carrying the nonpolar *ureI* deletion showed that UreI is essential for *H. pylori* survival in vivo and/or colonization of the mouse stomach. The replacement of *ureI* with the nonpolar cassette strongly reduced *H. pylori* survival in acidic conditions (1-h incubation in phosphate-buffered saline solution at pH 2.2) in the presence of 10 mM urea. UreI is predicted to be an integral membrane protein and may therefore be involved in a transport process essential for *H. pylori* survival in vivo.

*Helicobacter pylori* is a microaerophilic gram-negative bacterium which colonizes the gastric mucosa of humans (9). *H. pylori* is associated with gastritis and peptic ulcer disease and has been shown to increase the risk of gastric cancers. Urease is a major virulence factor of *H. pylori*. It is involved in neutralizing the acidic microenvironment of the bacterium and also plays a role in *H. pylori* metabolism (10, 24).

The urease-encoding region of the *H. pylori* genome is composed of two gene clusters common to all strains (8) (Fig. 1), one comprising the *ureAB* genes encoding the structural urease subunits and the other containing the *ureFGH* genes encoding the accessory proteins required for nickel incorporation into the urease active site. There is a gene of unknown function, *ureI*, immediately upstream from the latter gene cluster and transcribed in the same direction (Fig. 1). The distances separating *ureI* from *ureE* (1 bp) and *ureE* from *ureF* (11 bp) suggest that *ureI-ureE-ureF* constitutes an operon. Cotranscription of *ureI* and *ureE* has been demonstrated by Northern blot analysis (1). An *H. pylori* N6 mutant with the *ureI* gene disrupted by a MiniTn3-Km transposon was previously obtained (12). This strain (N6-*ureI*::TnKm-8) presented a urease-negative phenotype, so it was concluded that *ureI* was an accessory gene required for full urease activity.

The sequence of UreI from *H. pylori* and those of the AmiS proteins, encoded by the aliphatic amidase operons of *Pseudomonas aeruginosa* and *Rhodococcus* sp. strain R312, are similar (4, 25). Aliphatic amidases catalyze the intracellular hydrolysis of short-chain aliphatic amides to produce the corresponding organic acid and ammonia. We have shown that *H. pylori* also has such an aliphatic amidase, which hydrolyzes acetamide and propionamide in vitro (21).

The sequence similarity between UreI and AmiS together with the very similar structures of the urease and amidase substrates (urea:  $\text{NH}_2\text{-CO-NH}_2$ ; acetamide:  $\text{CH}_3\text{-CO-NH}_2$ ) and the fact that ammonia is produced by both enzymes

opened new perspectives for an investigation of the function of the *H. pylori* UreI protein.

**Construction of defined mutations of the *H. pylori* *ureI* gene.** *H. pylori* strains with defined mutations in *ureI* were generated by allelic exchange to determine whether the UreI protein was necessary for full urease activity. For this purpose, two plasmids (pILL823 and pILL834) with cassettes carrying antibiotic resistance genes inserted in *ureI* were constructed in *Escherichia coli*.

In one plasmid, pILL823 (Fig. 2), the *ureI* gene was inactivated by the insertion of a promoterless *cat* gene, conferring resistance to chloramphenicol (CM). A 780-bp blunt-ended *Bam*HI restriction fragment containing the "cat cartridge" from pCM4 (Pharmacia, Uppsala, Sweden) was introduced into a unique *Hpa*I site, between codons 21 and 22 of *ureI*, in pILL753 (8).

The second plasmid, pILL834, carried a *ureI* gene in which all but the first 21 codons were deleted and replaced with a nonpolar cassette (subcloned from pUC18K2 [18]) composed of the *aphA-3* kanamycin (KM) resistance gene (23) with its promoter and terminator regions deleted. In *Shigella flexneri* (18) and other organisms (such as *Yersinia enterocolitica* [2]), this cassette has been shown not to affect the transcription of the genes downstream within an operon as long as these distal genes have intact translation signals. There is only 1 bp separating *ureI* from *ureE* (Fig. 1), and *ureE* does not have a ribosome binding site (RBS) of its own; so the expression of *ureI* and *ureE* is transcriptionally and translationally coupled. Therefore, the *ureI* deletion was accompanied by the addition of an RBS immediately upstream from *ureE*. As shown in Fig. 2, three intermediates, pILL824, pILL825, and pILL833, were constructed in order to produce the final plasmid, pILL834.

**Introduction of *ureI* mutations into *H. pylori*.** *H. pylori* *ureI* mutants were produced by allelic exchange following electroporation with a concentrated preparation of pILL823 and pILL834 (as previously described [21]) of *H. pylori* N6 (11) and of mouse-adapted *H. pylori* SS1 (Sydney strain) (16). Bacteria showing chromosomal allelic exchange with pILL823 were selected on CM (4  $\mu\text{g/ml}$ ), and those with chromosomal allelic exchange with pILL834 were selected on KM (20  $\mu\text{g/ml}$ ). We

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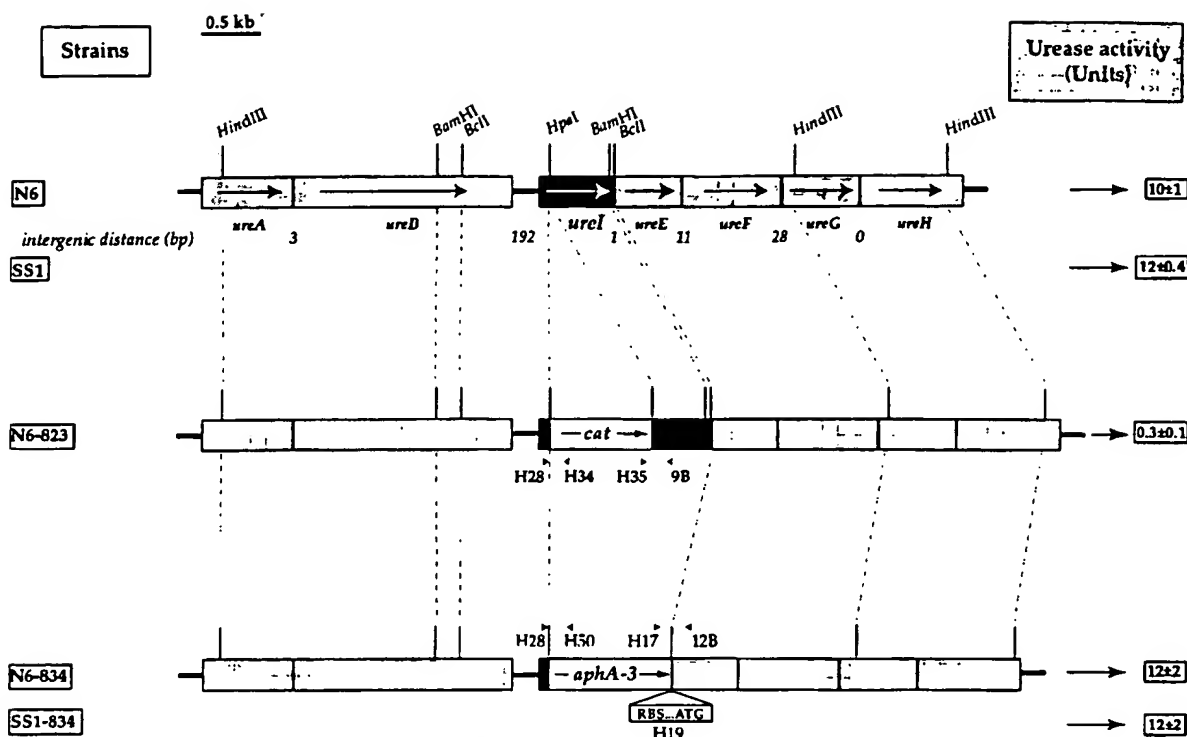


FIG. 1. The urease gene cluster of *H. pylori* parental strains N6 and SS1 and of the derived mutants deficient in *UreI*, strains N6-823, N6-834, and SS1-834. The genes are indicated by boxes with arrows showing the direction of their transcription. The distances between the *ure* genes are given in base pairs. The sites hybridizing to the primers used to confirm correct allelic exchange in strains N6-823, N6-834, and SS1-834 are shown. Blank boxes represent the cassettes containing the genes conferring resistance to CM (*cat*) or to KM (*aphA-3*). The urease activities of these strains are given on the right side of the figure. Urease activity was measured at pH7 as the release of ammonia in crude extracts of bacteria grown for 48 h on blood agar plates as described previously (8). One unit corresponds to the amount of enzyme required to hydrolyze 1  $\mu$ mol of urea/min/mg of total protein. The data are means  $\pm$  standard deviations calculated from three to five determinations.

checked that the desired allelic exchange had taken place in strains N6-823, N6-834, and SS1-834 (Fig. 1) by performing PCR with the appropriate oligonucleotides (Table 1). The PCR products obtained with genomic DNA of these strains were as expected: for strain N6-823, 140 bp with primers H28 and H34, 220 bp with primers H35 and 9B, and 1.2 kilobase pairs (kb) with primers H28 and 9B; for strains N6-834 and SS1-834, 150 bp with primers H28 and H50, 180 bp with primers H17 and 12B, and 1 kb with primers H28 and 12B.

The growth rate of strain N6-834 carrying mutant *ureI* with a nonpolar cassette was compared to that of the parental strain, N6. No difference in the colony size was observed on blood agar medium plates. Identical doubling times and stationary phase optical densities were measured for both strains grown in brain heart infusion (Oxoid) liquid medium containing 0.2%  $\beta$ -cyclodextrin (Sigma). *UreI* is thus not essential for *H. pylori* growth in vitro.

**Urease activities of *H. pylori ureI* mutants.** The urease activities of strains N6-823, N6-834, and SS1-834 were measured in vitro on crude extracts as described previously (8) and compared to the activities of the parental strains, N6 and SS1 (Fig. 1). Urease activity was almost completely abolished in strain N6-823 ( $0.3 \pm 0.1$  U). Strains N6-834 and SS1-834, with nonpolar *ureI* mutations, had wild-type levels of activity (N6-834 and SS1-834:  $12 \pm 2$  U; N6:  $10 \pm 1$  U; SS1:  $12 \pm 0.4$  U).

These results strongly suggested that the urease-negative phenotype of the N6-*ureI*::TnKm-8 strain (12) and the very weak urease activity of the N6-823 strain were due to a polar effect of the inserted cassettes on the expression of the downstream genes *ureE* and *ureF* (Fig. 1). This hypothesis was tested

by measuring the urease activity of strain N6-823 complemented in *trans* with an *E. coli/H. pylori* shuttle plasmid expressing the *ureEF* genes. This plasmid, pILL845 (Fig. 2), was obtained by insertion of a 2.8-kb *ClaiI*-*BamHI* fragment of pILL834 (comprising the 3' end of *ureB*, *ureI* with the nonpolar cassette replacing deleted codons, and intact *ureE* and *ureF* genes) into the corresponding sites of the pHel2 shuttle vector (14). Strain N6-823 was electroporated with a DNA preparation of pILL845 (as described in reference 21), and transformants were selected on KM (20  $\mu$ g/ml) and CM (4  $\mu$ g/ml). In strain N6-823 harboring pILL845, a high level of urease activity was restored (25 U), confirming that the very low level of urease activity of strain N6-823 was due to a polar effect on the expression of accessory genes *ureEF*.

**Colonization test for the *H. pylori* SS1-834 mutant in the mouse animal model.** The mouse model for infection by the *H. pylori* SS1 strain (Sydney strain) (16), validated in our laboratory (6, 13), was used to test the function of *UreI* in vivo. Mice were infected either with the nonpolar *ureI* mutant, SS1-834, or with the parental strain, SS1 (which had gone through an equivalent number of in vitro subcultures) as a positive control. This experiment was repeated three times and produced identical results. Two independently constructed SS1-834 mutants were used. The first mutant strain had gone through 30 in vitro subcultures; the second had gone through only 20. R. Ferrero (10a) showed that, under the same experimental conditions, strain SS1 can undergo up to 80 in vitro subcultures without losing its colonization capacity.

In each experiment, aliquots (100  $\mu$ l) containing  $10^6$  *H. pylori* SS1 or SS1-834 bacteria prepared in peptone broth were

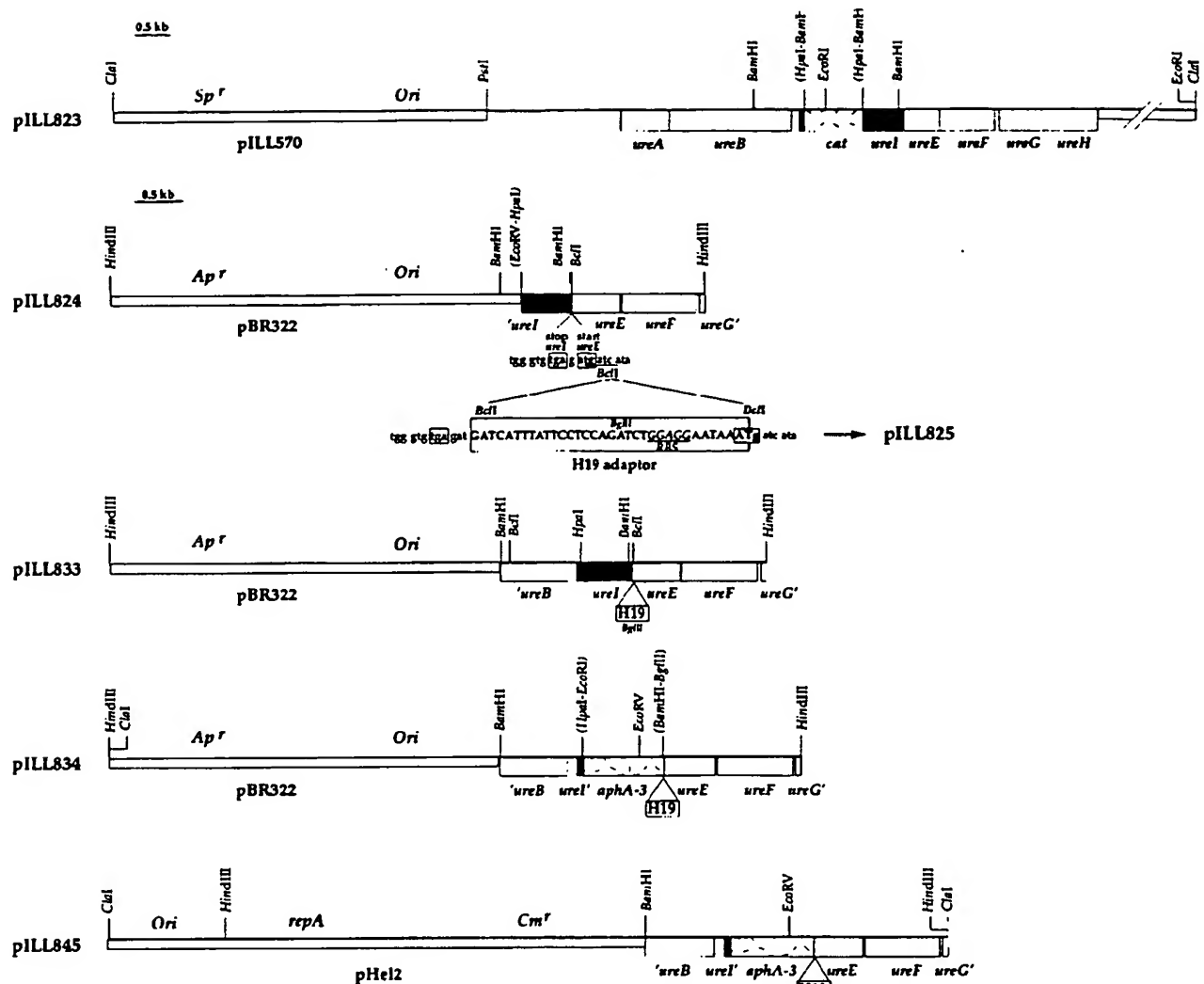


FIG. 2. Restriction map of pILL823, pILL824, pILL833, pILL834, and pILL845. Small boxes mark the vector of each plasmid; large boxes correspond to genes. *Ori* indicates the position of the ColE1 origin of replication. *repA* is the gene coding for the RepA protein, which is necessary for autonomous replication of pHel2 in *H. pylori*. *Sp<sup>r</sup>*, *Ap<sup>r</sup>*, and *Cm<sup>r</sup>* indicate the genes conferring resistance to spectinomycin, ampicillin, and CM, respectively. The sequence of the DNA region comprising the *ureI* stop codon and the *ureE* start codon, including the *BclI* site where adapter H19 was inserted, is given below pILL824, which was obtained by the insertion of a 1.8-kb *HpaI*-*HindIII* fragment from pILL753 (8) into pBR322. Plasmid pILL825 was produced by the insertion of the H19 adaptor (carrying an RBS and ATG in frame with *ureE*) (Table 1) into the *BclI* site of pILL824; the resulting *ureI-ureE* intergenic sequence is also shown. The stop codon of *ureI* and the start codon of *ureE* are boxed, and the RBS is underlined. In pILL833, the *BamHI* fragment of pILL825 was replaced by a 1.3-kb blunt-ended *PvuII*-*BamHI* fragment from pILL753. Plasmid pILL834 was obtained by replacement of the *HpaI*-*BglII* fragment of pILL833 with an 850-bp blunt-ended *EcoRI*-*BamHI* fragment of pUC18K2 containing the nonpolar KM cassette (18). Parentheses indicate the position of restriction sites removed by ligation.

administered orally to 10 mice each (6- to 8-week-old Swiss specific-pathogen-free mice) as described by Ferrero et al. (13). Mice were killed 4 weeks after inoculation. We tested for the presence of *H. pylori* with a direct urease test on biopsies performed on half the stomach (13). The remaining gastric tissues were used for quantitative culture of *H. pylori* as described by Ferrero et al. (13). In every experiment, the stomachs of the 10 SS1-infected mice all tested positive for urease. The bacterial load was between  $5 \times 10^4$  and  $5 \times 10^5$  CFU per g of stomach tissue. None of the stomachs of the mice infected with strain SS1-834 tested positive for urease, and no *H. pylori* cells were cultured from them. Thus, the UreI protein is essential for *H. pylori* in vivo survival and/or colonization of the mouse stomach.

TABLE 1. Names and nucleotide sequences of oligonucleotides used in this study

Primer <sup>a</sup>	Oligodeoxynucleotide sequence (5' to 3')
H17	TTTGACTTACTGGGATCAAGCCTG
H19	GATCATTTATTCCTCCAGATCTGGAGGAATAAAT
H28	GAAGATCTCTAGGACTTGATTGTTATAT
H34	TATCAACGGTGGTATATCCAGTG
H35	GCAGTTATTGGTGCCTTAAACG
H50	CCGGTGATATTCTCATTTTAGCC
8A	GCGAGTATGTAGTTTCAGTA
9B	GTGATACCTGAGCAATATCTTCAGC
12B	CAAATCCACATAATCCACGCTGAAATC

<sup>a</sup> H19 was used as the adaptor, and the others were used as primers for PCR amplification.

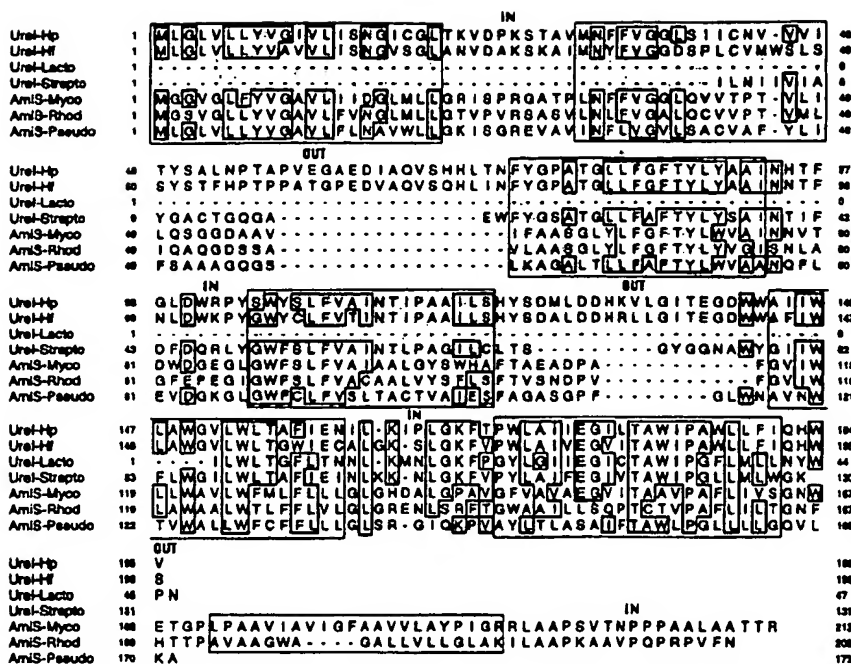


FIG. 3. Alignment of the amino acid sequence of Urel from *H. pylori* with those of similar proteins and prediction of the two-dimensional structure of members of the protein family comprising Urel and AmiS. Residues identical at one position in at least four sequences are boxed; dashes indicate gaps inserted to optimize alignment. The organisms from which the sequences originated and the degree of identity of each with the *H. pylori* Urel protein are as follows: Urel-Hp, *H. pylori* (195 residues; accession no., M84338); Urel-Hf, *H. felis*; 74% identity over 196 residues (accession no., A41012); Urel-Lacto, *L. fermentum*; 55% identity over the 46-residue partial sequence (accession no., D10605); Urel-Strepto, *S. salivarius*; 54% identity over the 129-residue partial sequence (accession no., U35248); AmiS-Mycos, *M. smegmatis*; 39% identity over 172 residues (accession no., X57175); AmiS-Rhod, *Rhodococcus* sp. strain R312; 37% identity over 172 residues (accession no., Z46523); and AmiS-Pseudo, *P. aeruginosa*; 37% identity over 171 residues (accession no., X77161). Predicted transmembrane  $\alpha$ -helices are shown as shaded boxes. The regions separating these boxes are hydrophilic loops labeled IN when they are predicted to be intracellular and OUT when they are predicted to be extracellular.

**Survival of the *H. pylori* N6-834 mutant in acidic conditions.** Survival under acidic conditions in the presence of 10 mM urea or in the absence of urea was tested with strains N6 and N6-834. The experimental procedures were those described by Clyne et al. (7). Exponentially grown bacteria were harvested and washed in phosphate-buffered saline (PBS; Boehringer, Mannheim, Germany), and  $2 \times 10^8$  CFU of bacteria per ml were resuspended in PBS at pH 2.2 or 7 in the presence of 10 mM urea or in the absence of urea and incubated for 1 h at 37°C. To evaluate bacterial survival, quantitative cultures (5 days of growth) of the *H. pylori* strains were performed. At pH 7, in the absence of urea, both strains survived similarly ( $10^8$  CFU/ml). In agreement with the results of Clyne et al. (7), none of the strains survived at pH 7 in the presence of urea because the final pH rose to 9. As expected (7), both strains were killed at pH 2.2 in the absence of urea, and significant survival at pH 2.2 in the presence of urea was observed with strain N6 ( $5 \times 10^6$  CFU/ml; final pH 6.5). In contrast, when the nonpolar *ureI* mutant strain N6-834 was incubated at pH 2.2 in the presence of urea, a low level of survival was observed ( $10^3$  CFU/ml) and the pH was unchanged (pH 2.3) after 1 h of incubation.

**Alignment of the Urel and AmiS protein sequences and two-dimensional structure prediction.** A systematic search for Urel homologs in the protein data banks was carried out. We found that *H. pylori* is not the only ureolytic bacterium with a *ureI* gene. Two phylogenetically related gram-positive organisms, *Streptococcus salivarius*, a dental plaque bacterium (5), and *Lactobacillus fermentum*, a lactic acid bacterium (15), carry genes (the available sequences are only partial) coding for Urel homologs (Fig. 3) located immediately upstream from the ure-

ase structural genes. The *ureI* gene has also been detected in various *Helicobacter* species; the *Helicobacter felis ureI* gene has been entirely sequenced (10b) (Fig. 3). PCR experiments have suggested that there is a *ureI* gene in *Helicobacter heilmannii* (22) and in *Helicobacter mustelae* (unpublished data).

Sequence similarities between the Urel protein of *H. pylori* (21.7 kDa) and the AmiS proteins expressed by the aliphatic amidase operons from *P. aeruginosa* (25) and *Rhodococcus* sp. strain R312 (4) have been reported. In *Mycobacterium smegmatis*, there is an additional AmiS homolog encoded by a gene, open reading frame P3, located immediately upstream from an amidase gene (17).

Alignment of these Urel and AmiS proteins [with the Clustal W(1.60) program] defined strongly conserved stretches of amino acids (Fig. 3). All but one of these conserved blocks are in highly hydrophobic segments. These regions, each 17 to 22 residues long, are probably folded into transmembrane  $\alpha$ -helices (Fig. 3). Six transmembrane regions were predicted for the proteins from *H. pylori*, *H. felis*, and *P. aeruginosa*, and seven were predicted for those from *Rhodococcus* sp. strain R312 and *M. smegmatis* (these are highly reliable predictions, performed with pH-D, a profile-fed neural network system [20]). The orientations of the Urel and AmiS proteins in the membrane were deduced (20) from the charges of the intercalated hydrophilic regions, which are short in these proteins (Fig. 3). These results strongly suggest that the members of the family comprising Urel and AmiS, found in both gram-positive and -negative bacteria, are integral membrane proteins. These proteins have no signal sequence and should therefore be inserted into the cytoplasmic membrane in gram-negative bacteria.



**Conclusions.** The urease cluster of *H. pylori* is unique among the many urease operons of gram-negative bacteria that have been sequenced (19) in that it has an extra gene, *ureI*. The function of *UreI* has therefore been the subject of much speculation. It has mostly been assigned the function of an accessory protein required for nickel incorporation at the urease active site or of a nickel transporter. We have demonstrated that *UreI* is not required for full activation of *H. pylori* urease during in vitro growth. *UreI* is thus not a nickel transporter since such a protein, NixA (3), already identified in *H. pylori*, is necessary for full urease activity. We showed herein that replacing *ureI* with a nonpolar cassette has no effect on urease activity measured in vitro. This is the first time that a nonpolar cassette (18) has been shown to be functional in *H. pylori*. This will certainly be a valuable tool for genetic analysis of complex *H. pylori* operons.

We observed that *UreI* was essential for survival in vivo and/or for colonization of the mouse stomach. This could be due to the reduced resistance to acidity of the *ureI* mutant, as suggested by the results of tests of in vitro survival in acidic conditions with 10 mM urea. *UreI* has a sequence similar to those of the AmiS proteins, proposed to be involved in the transport of short-chain amides (25), molecules structurally similar to urea. The *UreI* and AmiS proteins have the characteristics of integral membrane proteins, probably of the cytoplasmic membrane. Different roles for *UreI* can tentatively be proposed. *UreI* might be involved in (i) transport of urea or short-chain amides, (ii) an uptake system for maintaining appropriate intracellular ammonia concentrations, or (iii) the export of excess intracellular ammonium. An essential role for *UreI* as an amide transporter seems unlikely because in mouse colonization experiments (performed as described above) an SS1 mutant deficient in aliphatic amidase (carrying the mutation described in reference 21) colonized mice as efficiently as the parental strain, SS1. In addition, amidase activity was not significantly modified by the deletion of *ureI* in strain N6-834. Our results concerning acidity survival are most compatible with *UreI* being involved in ammonium export.

Finally, *UreI*, as a membrane protein essential for the survival of *H. pylori* in vivo, is an interesting potential target for new antibacterial drugs.

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